(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 15 February 2001 (15.02.2001)

PCT

(10) International Publication Number WO 01/10459 A2

- (51) International Patent Classification7:
- A61K 39/00
- (21) International Application Number: PCT/GB00/02973
- (22) International Filing Date: 7 August 2000 (07.08.2000)
- (25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

9918591.0 9923030.2 7 August 1999 (07.08.1999) GB 30 September 1999 (30.09.1999) GB

- (71) Applicant (for all designated States except US): AQUA HEALTH (EUROPE) LIMITED [GB/GB]: Enterprise House, Springkerse Business Park, Stirling FK7 7UF (GB).
- (72) Inventor; and
- (75) Inventor/Applicant (for US only): BARNES, Andrew, Cartner [GB/GB]; 54 Union Grove, Aberdeen AB106RX (GB).

- (74) Agent: MURGITROYD & COMPANY: 373 Scotland Street, Glasgow G5 8QA (GB).
- (81) Designated States (national): AE. AL. AM. AT. AU. AZ. BA. BB. BG. BR. BY. CA. CH. CN, CR. CU. CZ. DE. DK. DM. EE. ES. FI. GB. GD. GE. GH. GM. HR. HU. ID. IL. IN. IS. JP. KE, KG. KP. KR. KZ. LC. LK. LR. LS. LT. LU. LV. MA. MD. MG. MK. MN. MW. MX. NO. NZ. PL. PT. RO. RU. SD. SE, SG. SI. SK, SL, TJ, TM. TR. TT. TZ. UA. UG. US. UZ. VN. YU. ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

 Without international search report and to be republished upon receipt of that report.

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

A2

(54) Title: VACCINE

(57) Abstract: A vaccine composition is provided for the prophylactic and/or therapeutic treatment of fish for infection by bacteria, particularly by the organism *Photobacterium damselae* subsp. *piscicida*. The composition comprises components produced by a culture of the organism. The culture is treated to kill the organism prior to use, preferably by treatment with formalin after components have been produced. Two of the components, involved in invasion of host cells and produced in high quantities when the culture is grown in medium containing excess iron, induce production of antibodies on injection into fish which prevent the entry of the organism into fish cells. These antibodies protect the fish from infection by the organism.

1

1	"Vaccine"
2	
3	The present invention relates to novel compositions
4	for use as vaccines and medicaments for the
5	prophylactic treatment of fish for infection by
6	bacterial organisms, and particularly for the
7	protection of fish from infection by bacteria such as
8	Photobacterium damselae subsp. piscicida.
9	
10	Photobacterium damselae subsp piscicida, formerly
11	Photobacterium damsela subsp. piscicida was
12	reclassified from, and formerly reported as
13	Pasteurella piscicida and reallocated to the family
14	vibrionaceae based on small subunit 16s ribosomal RNA
15	homology (Gaulthier et al (1995) Int. J. Syst.
16	Bacteriol. 45:139-144). P. damselae causes serious
17	fatal disease, commonly called pseudotuberculosis, in
18	warm water marine fish infecting major organs and
19	tissues resulting in characteristic lesions or
20	pseudotubercles in the musculature.
21	
22	Appearance of antibiotic resistance coupled with the
23	observation that fish cease feeding when infected,

60: 2990-2998). These include siderophores and iron 32 (Magarinos et al. (1994) Appl. Environ. Microbiol. ₽£ high affinity iron uptake systems are expressed 33 agents which reduce its avilability, a variety of 35 sufficient in the medium, or by including chelating Ţξ which iron is limiting, either by not including 30 exception. It has been shown that under conditions in 62 absolute requirement for iron, and P. damselae is no 82 It is an established fact that bacteria have an 72 97 in commercial situations. 52 Pathol.14:120-122) but these have not been effective ₽2 (Magarinos et al. (1994) Bull. Eur. Ass. Fish 23 extracellular products (ECP) have been described 22 proteins (IROMPs) and bacterins enriched with ZZ Preparations based on iron restricted outer membrane 02 antigens contained in these preparations. 6 T Furthermore, little is known of the protective 8 T et al. (1997) Fish Shellfish Immunol. 8: 63-72). LI response of the host to the infectious agent (Arijo 9 T ppl67-177). This may reflect the low antibody SI (1997) Fish Vaccinology, Karger, Basel, Switzerland, ÐΤ reproducible protective efficacy (Romalde & Magarinos ετ however, many of these have failed to yield IS based on formalin killed bacteria (bacterins), ττ Various vaccination strategies have been suggested οτ 6 means of controlling this disease. 8 through immunisation appears to be the most desirable L Microbiology 145: 483-494). Therefore prophylaxis 9 during host colonisation (Barnes et al. (1999) S evidence to suggest a phase of intracellular survival ₽. failure. Furthermore, there is an increasing body of έ have been suggested as reasons for chemotherapeutic Z and thus do not consume antibiotic medicated feeds

3

restricted outer membrane proteins (IROMPS). In other 1 bacterial species such as Aeromonas salmonicida 2 (Hirst & Ellis (1994) Fish Shellfish Immunol. 4:29-3 45) and Pasteurella haemolytica (UK patent 4 specification 8805253 W Donachie, UK) IROMPs have 5 been exploited as protective antigens in successful 6 vaccines. However these do not appear to be 7 protective in P. damselae, (Romalde & Magari-os 8 (1997) Fish Vaccinology, Karger, Basel, Switzerland, 9 pp167-177). This lack of protection may reflect the 10 strategy for infection by P. damselae. Under iron 11 limitation, P. damselae produces increased protease 12 activity (Bakopolous (1997) J. Fish Dis. 20: 297-305) 13 and increased capsular polysaccharide. Capsular 14 polysaccharide increases resistance to serum killing 15 and prevents phagocytosis by host macrophages (Arijo 16 et al. (1998) Fish Shellfish Immunol. 8: 63-72.). 17 This is significant as P. damselae is not resistant 18 to attack by phagocytes (Skarmeta et al. (1995) Dis. 19 Aquat. Org. 23:51-57) as a result of its lack of 20 adaptive response to killing by reactive oxygen 21 species produced during the macrophage respiratory 22 burst (Barnes et al. (1999) Microbiology 145: 483-23 495). However, the increased proteolytic activity is 24 able to free iron in the form of haem through red 25 blood cell lysis. Haem is absorbed by the capsule 26 (Ana Do Vale, Pers. Comm.) and the iron translocated 27 across the membrane by the IROMPs. Thus with capsular 28 polysaccharide acting as the first level of iron 29 uptake the IROMPs are not exposed during the disease 30 process. As iron levels increase, capsular production 31 decreases, IROMPs are switched off and proteolytic 32 activity becomes undetectable (Bakopolous (1997) J. 33 Fish Dis. 20: 297-305). This would leave P. damselae 34 susceptible to both antibody and macrophage attack. 35

WO 01/10459

However, at this point the organism is able to adhere 2 to (Yoshida et al. (1997) J. Fish Dis. 20: 77-80) and enter non-phagocytic fish cells (Magari-os et al. 3 4 (1996) FEMS Microbiology Lett. 138: 29-34) thus 5 avoiding attack by antibodies or phagocytes. 6 7 It is an aim of the present invention to provide a vaccine for the protection of fish from infection by 8 9 bacteria such as Photobacterium damselae. 10 It is another aim of the invention to provide a 11 12 process for the production of such a vaccine. 13 According to the present invention there is provided 14 a vaccine and/or therapeutic composition comprising 15 biological material derived from a culture of 16 17 Photobacterium damselae, characterised in that the 18 bacterial cells have been cultured in a culture 19 medium containing excess iron over that which is 20 required for normal growth of the bacterium. 21 By "containing excess iron" the culture medium should 22 23 contain at least twice the amount of iron in standard tryptone soya broth (Oxoid). 24 25 Preferably the culture medium contains between $0.5\mu M$ 26 27 and 1mM iron. 28 29 More preferably the medium contains between $10\mu M$ and 30 $500\mu M$ iron. 31 32 Most preferably the medium contains between $25\mu M$ and 33 500µM iron. 34

5

1	Typically an outermembrane protein which may be						
2	involved in entry into host cells or invasin or						
3	adhesin is expressed at levels higher than those in						
4	normal culture medium.						
5							
6	By "expressed at levels higher than those in normal						
7	culture medium" the protein or invasin or adhesin is						
8	expressed at at least twice the normal expression						
9	levels. Normal expression levels are levels of						
10	expression in standard tryptone soya broth (Oxoid).						
11							
12	Typically an extracellular protein serologically						
13	related to invasin is expressed at higher levels than						
14	in normal culture.						
15							
16	Typically a 55Kda extracellular protein complex						
17	and/or a 97Kda outermembrane protein is expressed at						
18	levels higher than those in normal culture medium.						
19							
20	The invention also provides purified proteins as						
21	described for use as vaccines or in the preparation						
22	of vaccines.						
23							
24	The invention also provides antibodies to the						
25	purified proteins.						
26							
27	The invention further provides a method for						
28	production wherein the cells are cultured in a						
29	culture medium containing sufficient iron such that						
30	the cells are loaded up with iron to the point where						
31	linear uptake of iron from the culture medium no						
32	longer occurs.						
33							
34	Preferably the cells are saturated with iron.						
35							

6

1 Preferably iron is supplied to the medium in the form 2 of a ferric salt. 3 4 Preferably the bacteria have been inactivated after 5 culture. 6 7 Preferably the inactivation is carried out using a 8 formaldehyde composition. 9 The invention further provides cells or cell 10 11 membranes or extracellular products of the cultured 12 bacteria. 13 14 The biological material is preferably provided in a 15 physiologically acceptable carrier. 16 17 The composition of the invention preferably includes an adjuvant suitable for enhancing immunological 18 19 response. 20 21 A vaccine composition is thus provided for the 22 prophylactic and/or therapeutic treatment of fish for 23 infection by bacteria, particularly by the organism 24 Photobacterium damelae subsp. piscicida. 25 26 The composition comprises components produced by a 27 culture of the organism. The culture is treated to kill the organism prior to use, preferably by 28 treatment with formalin after components have been 29 30 produced. 31 32 Two of the components, involved in invasion of host 33 cells and produced in high quantities when the culture is grown in medium containing excess iron, 34 induce production of antibodies on injection into 35

1

7

fish which prevent the entry of the organism into 2 fish cells. 3 These antibodies protect the fish from infection by 4 5 the organism. 6 7 The invention thus also provides antibodies. 8 9 In one particular embodiment the biological material 10 is derived from the culture of Photobacterium damselae strain MT1415 deposited under Accession No 11 12 41062 on 4 August 2000 at N.C.I.M.B. in Aberdeen, 13 United Kingdom. 14 15 The present inventor has determined that a protein 16 (invasin or adhesin) expressed in the outer membrane 17 under iron replete conditions is involved with internalisation of P. damselae by Sea Bass 18 19 Fibroblastic Cells. Lectins which specifically bind this protein inhibit the invasive capacity of P. 20 21 damselae, whilst lectins which do not bind the 22 protein do not significantly inhibit invasion of Sea Bass Larval fibroblast cell lines (SBLs). 23 Furthermore, antibodies raised against this protein 24 25 in sea bass prevent internalisation of the bacterium 26 by fish epithelial cells (EPCs). The expression of 27 this protein can be increased by addition of ferric iron in the form of ferric chloride, to the growth 28 29 For example, addition of one hundred micromolar (100 μM) ferric chloride increases 30 expression of the protein by at least 2.6 fold. 31 32 Bacterins produced by growing P. damselae under such iron supplementation are protective against clinical 33 34 pasteurellosis in rainbow trout and juvenile sea 35 bream.

8

1 2 Iron may be supplied to the culture medium in any 3 form that results in its uptake by the bacterial cells in such a manner that increase of the invasin 4 5 results. Other culture conditions may also increase expression of this protein. For example more may be 6 7 expressed under anaerobiosis in the presence of sufficient iron. However, higher cell densities are 8 9 achieved by growing the bacteria between 22°C and 35°C with continous shaking, or other aeration, to 10 11 maintain the oxygen content. 12 The vaccine cells of the invention are inactivated by 13 any standard method, but conveniently by use of 14 15 formaldehyde. Further preparation of the vaccine such as addition of an adjuvant, concentration of the 16 cells, or resuspension into an acceptable carrier may 17 afford additional protection. 18 19 The vaccines of the invention and a method for their 20 21 production will now be described by way of illustration only by reference to the following non-22 limiting Examples and Figures. Further embodiments 23 falling within the scope of the claims will occur to 24 those skilled in the art in light of these. 25 26 27 Figures 28 Figure 1. Shows a histogram of relative percent 29 survival (RPS) vs vaccine treatment for rainbow trout 30 challenged with organisms of genus Photobacterium. 31 32 Figure 2. Shows a histogram of RPS vs vaccine 33 treatment for gilthead seabream (Sparus aurata) 34 challenged with organisms of genus Photobacterium. 35

WO 01/10459

1 2 Figure 3. Shows a Western blot of P. damselae outer 3 membrane proteins (OMP, lanes 2,4,6,8) and 4 extracellular products (ECP, lanes 3,5,7,9) stained 5 with aurodye (for protein, lanes 1-5) and sea bass vs iron-supplemented P. damselae (lanes 6-9), Clearly 6 7 showing 97KDa protein (lane 8) and Ô55Kda protein 8 complexÕ (lanes 7, 9). 9 10 Figure 4. Shows percentage of EPC cells with at least one intracellular P. damselae preincubated with 2) 11 12 normal sea bass serum or 3) sea bass vs P.damselae invasin antiserum with 1) control. 13 14 15 Figure 5. Shows invasion of EPC/SBL cells by 16 P.damselae incubated with various lectins, relative to controls not previously incubated with lectins. 17 18 Figure 6. Shows Western blot of P.damselae OMPs 19 20 produced under iron supplemented conditions, probed 21 with Lectins and sea bass vs P. damselae antiserum and 22 rabbit anti 55KDa complex.Lanes: 1) rabbit vs 55KDa 23 complex antibody; 2) bass vs MT1415 antibody 3) 24 biotinylated dolichos biflorus agglutinin; 4) biotinylated Concanavalin A lectin; 5) Sigma 25 26 biotinylated molecular weight markers. 27 28 Figure 7 shows Western blot, stained with Sea Bass vs Photobacterium damselae antibody, of Outer membrane 29 poteins from vaccine preparations showing absence of 30 31 55KDa complex in iron depleted culture (Vaccine B), 32 compared to iron supplemented culture (Vaccine A). 33 34 Figure 8 shows relative percent survival (RPS) of Sea 35 bass Dicentrarclus Labrax challenged with

10

Photobacterium damselae subsp. piscicida following 1 2 vaccination with either A) Vaccine expressing 55KDa 3 complex and 97KDa OMP or B) Vaccine NOT expressing 4 55KDa complex or 97KDa OMP (see also Figure 7). 5 6 Example 1: Production of an invasin-expressing P. 7 damselae vaccine 8 9 Tryptone soya broth (Oxoid) was made up in distilled water at 30g/l and sodium chloride was added at 20g/l 10 in conical flasks such that the volume of the flask 11 was five times the volume of the medium to allow for 12 sufficent aeration. After sterilisation by 13 autoclaving at 121 °C for 15 minutes, ferric chloride 14 was added to a final concentration of 100 micromolar 15 from a sterile stock of 100 millimolar in distilled 16 17 water. The broth was pre warmed to 25 °C and seeded with 1/10000th volume of an overnight tryptone soya 18 broth culture, containing 2% salt, of Photobacterium 19 20 damselae (strain MT1415, capsule positive virulent 21 isolate as deposited under Accession No 41062 on 4 22 August 2000 at N.C.I.M.B in Aberdeen, United 23 Kingdom). Incubation was continued at 25 °C with shaking at 140 rpm for 40 hours. After incubation, 24 the culture was inactivated by addition of 0.5% 25 26 formalin v/v (0.2% free formaldehyde) and the broth was left at 25 °C for 24 hours to allow complete 27 inactivation. The optical density of the final 28 vaccine preparation was determined and the vaccine 29 30 was stored at 4 °C until required. 31 32 For administration this cell suspension was 33 administered in one of a number of ways:

34

11

1 Rainbow trout (10-15g) held in freshwater at 25 °C 2 were anaesthetised with MS222 (sigma). Anaesthetised fish were vaccinated by intraperitoneal injection of 3 neat vaccine preparation (100 μ l) or reference 4 preparation or sterile phosphate buffered saline as a 5 control. At least 500 degree days post immunisation, 6 fish were challenged by injection of 108 cfu of 7 virulent P. damselae intraperitoneally. Mortalities 8 were recorded daily and the RPS (relative percent 9 10 survival) as compared to control fish was determined. Results are shown in Figure 1 below wherein the RPS 11 using the present invention is compared with a number 12 of reference vaccines. 1, reference vaccine; 2, 13 reference vaccine; 3, Iron-restricted vaccine; 4, 14 iron supplemented vaccine (vaccine of the invention); 15 16 5, standard tryptone soya broth (TSB) vaccine; 6-12, reference vaccines. 17 18 Alternatively, sea bream, 0.3 g were immunised by 19 20 single 60 second immersion in a tenfold dilution of 21 the present invetion or reference vaccine. 500 degree days post vaccination fish were challenged by 22 immersion in a suspension containing approximately 23 10⁵ cfu/ml virulent P. damselae for 1 hour at 25 °C. 24 Mortalities were recorded as described above. Results 25 are given in Figure 2: 1, iron restricted vaccine; 2, 26 TSB vaccine; 3, Iron-supplemented vaccine 27 28 (invention). 29 30 Identification protective antigens 31 Vaccine prepared as described above was used to 32 immunise sea bass (Dicentrarchus labrax) by 33 intraperitoneal injection. Freunds incomplete 34 35 adjuvant was administered simultaneously at a ratio

12

of 1:1. Fish were maintained in seawater at 25 °C for 1 2 20 days after which they received a second identical 3 dose. After a further 20 days fish were bled and sera collected by allowing the blood to clot and removing the red blood cells by centrifugation. Sera were 5 6 dialysed against phosphate buffered saline and stored frozen at -80 °C. 7 8 9 Outer membrane proteins (OMP) and extracellular 10 products (ECP) were prepared from P. damselae grown 11 under iron limitation, in standard TSB, or under iron 12 supplementation. For iron limitation, 2,2 dipyridyl (100 micromolar) was added to tryptone soya broth to 13 chelate iron prior to inoculation with P. damselae. 14 15 Incubation was then carried out as described 16 above.OMP was prepared by precipitaion following sarkosyl solubilisation of the inner membrane 17 (Hancock & Poxton (1988) Bacterial Cell Surface 18 19 Techniques, John Wiley & Sons, Chichester, UK). Extracellular products were recovered from broth 20 21 culture supernatants. 22 23 ECPs and OMP (equal protein concentrations) were run 24 on SDS-PAGE gels under non-reducing conditions and 25 blotted onto PVDF membrane. Membranes were probed with bass vs iron-supplemented P. damselae, followed 26 by mouse vs bass immunoglobulin monoclonal antibody 27 previously described (Santos et al. (1997) Fish. 28 Shellfish Immunol.7:175-191), followed by goat vs 29 30 mouse conjugated with an alkaline phosphatase enzyme. To visulalise bands, membranes were incubated in 31 substrate consisting of Nitroblue tetrazoleum and 5-32 Bromo-4-chloro-3-indolyl phosphate. The results are 33 shown in figure 3 and figure 7. In the OMP-34 preparations from iron containing cultures a clear 35

13

1 band is visible close to the 97Kda marker which is 2 not present under iron limitation (figures 3 and 7). 3 By scanning densitometry this band was determined to 4 be at least 2.6 fold more concentrated in OMPs from 5 iron-supplemented cultures than from standard TSB 6 cultures, and not detectable in iron limited 7 cultures. In ECPs from cells grown in iron containing media, two bands were evident, one running close to 8 9 97Kda appeared to be the same as the band seen in OMP 10 preparations. The other was smaller, running close 11 to, but below, the 55KDa marker. Neither of these 12 bands were detected in OMPS or ECP from cells cultured in iron deficient media. 13 14 15 Inhibition of invasion of fish epithelial cells by antisera raised against iron-supplemented P. damselae 16 17 The ability of P. damselae to invade fish epithelial 18 19 cells (EPC) was determined by a fluorescent labelling 20 direct count method described by Bandin et al. (1995) 21 Dis Aquat. Org. 23: 221-227. P. damselae cells were 22 labelled using fluorescein isothiocyanate 23 (FITC) (0.1mg/ml) for 1 hour. P. damselae were then washed extensively in PBS and resuspended to a 24 density of 10^9 cells /ml. The cell suspension was 25 26 split and an aliquot incubated with heat inactivated 27 immune serum prepared by heating sea bass vs iron-

supplemented P. damselae antisera described above at 45 °C for 15 minutes, whilst a second aliquot was

P.damselae cells were added to 6 \times 10⁵ EPC cells in

1ml G-MEM, and allowed to attach and invade for 2

hours at 25 °C. External bacteria were removed by washing in PBS and EPCs were counter-stained with

incubated with heat-inactivated normal sea bass serum. Aliquots (10 microlitres) of serum treated

28

29 30

31

32

33 34

35

14

ethidium bromide. Aliquots (10 μ l) were placed on 1 2 glass slides and covered with a coverslip before analysis by fluorescent microscopy. Proportions of 3 4 internalised bacteria were determined by direct 5 counting at least 100 fields. Experiments were 6 replicated four times. The results are presented in 7 figure 4: 1, controls, P. damselae with no serum; 2, P. damselae incubated with heat inactivated normal 8 9 bass serum; 3, P. damselae incubated with heat inactivated antiserum (bass vs P. damselae) 10 11 12 Identification of the protein associated with P. damselae internalisation in Sea Bass Cells using 13 14 lectins 15 Previous work suggested that the entities involved in 16 17 internalisation of P. damselae in fish cells may be glycoproteins (Magari-os et al.(1996) FEMS Microbiol 18 Lett. 138: 29-34). The carbohydrate side chains of 19 20 glycoproteins can be specifically bound by certain 21 lectins. Lectins are plant extracts with highly 22 specific affinities for configurations of certain sugars. Incubating P. damselae with different lectins 23 then determining its ability to invade sea bass 24 larval fibroblast cells (SBL) identified lectins 25 which were capable of inhibiting invasion and those 26 which were not. The glycoproteins involved in 27 28 invasion could then be identified by probing Western blots with biotinylated lectins. 29 30 P. damselae, labelled with FITC, washed and 31 resuspended to a density of 109 cfu/ml as described 32 above, was incubated for 1 hour with various lectins 33 34 (Vector Laboratories) at a concentration of 100µg 35 lectin/ml. After incubation, the cells were washed

....

15

1 extensively and invasion assays carried out as described above. The results are presented in figure 2 5: P.damselae incubated with: 1, Sophora japonica 3 agglutinin; 2, Concanavalin A agglutinin; 3, Lens 4 5 culinaris agglutinin; 4, Griffonia simplicifolia 6 agglutinin; 5, succinylated wheatgerm agglutinin; 6, 7 Dolichos biflorus agglutinin; 7, peanut agglutinin; 8, soybean agglutinin; 9, Ulex europaeus agglutinin; 8 10, wheatgerm agglutinin. 9 10 Two lectins which were able to strongly inhibit 11 invasion by P. damselae, Sophora japonica agglutinin 12 13 (SJA) and ConA agglutinin (ConA) were selected. One lectin which did not inhibit invasion, Dolichos 14 biflorus agglutinin (DBA) was also selected. 15 Biotinylated preparations of these lectins (Vector 16 17 Laboratories) were used to stain Western blots of 18 SDS-PAGE-separated OMPs from P. damselae. results are presented in figure 6: 19 20 21 All the lectins stained a number of carboydrates/glycoproteins. However, only one region 22 was stained ConA, but not stained by DBA. This region 23 consisted of a complex of three protein bands and had 24 an approximate molecular weight of 55Kda under non 25 26 reducing conditions and was only detected in OMP preps from P. damselae cultured under iron-replete 27 conditions, not in preparations for P. damselae 28 cultured under iron limitation. Furthermore, when 29 Western blots of OMPs from P.damselae were cut and 30 stained with both ConA lectin and sea bass vs iron-31 32 supplemented P. damselae antiserum a common band, the 33 55 Kda complex, was stained by both methods.Lane 1 34 shows OMPs probed with rabbit vs 55KDa complex, Lane 2 shows OMPs probedwith bass vs Photobacterium 35

WO 01/10459

16 1 damseale MT1415 antiserum used in invasio0n 2 inhibition study. Lane 3 shows OMPS probed with 3 dolichos biflorus agglutinin, Lane 4 shows OMPs 4 probed with Concanavalin A lectin, Lane 5 shows Sigma 5 biotinylated molecular weight markers (SDS-6B). 6 7 Serological relationship between the 97 KDa OMP and the 55 KDa ECP. 8 9 10 The 97 KDa OMP and the 55KDa ECP were carefully 11 excised from polyacrylamide gels, homogenised and 12 injected into sea bass. After 30 days, sera were collected and used to probe Western blots of OMPs and 13 ECPs. Antisera raised in sea bass against the 97Kda 14 15 OMP cross reacted with the 55Kda ECP. Similarly, Antibodies raised in sea bass against the 55Kda ECP 16 also cross reacted with the 97KDa OMP. The inventor 17 suggests that the 55Kda ECP is a secreted version of 18 19 the 97Kda OMP. 20 21 Purification and N-terminal sequence 23 Subsequent purification of the 55Kda protein and

22

24 sequencing have revealed three proteins in this 25 region. The major antigenic protein is N-terminal blocked, consistant with glycosylation post 26 27 transcription, and therefore unable to obtain a sequence, however this fraction has strong 28 29 Haemagglutinating activity, suggesting probable 30 involvement in internalisation. A second protein gave 31 an N-terminal amino acid sequence with 100% homology to B-1,4 N-acetyl muramidase, a defence against other 32 bacteria: AMKRHGLDNYRGYSLGNWVC. 33 34 The third protein may be a fragment of a deaminase or 35 catabolic dehydratase: NVVLHGDNFDSTXVXVKAV.

17

1 2 3 Comparison of protective efficacy of vaccines which 4 5 express the 55KDa complex with vaccines which do not in a challenge study in Sea Bass (Dicentrarchus 6 7 labrax). 8 The following study was performed independently at 9 10 CEFAS Weymouth Laboratory under study protocol P0075, reference 99008). 11 Vaccines expressing the 55KDa protein complex were 12 prepared as follows: 500 ml Tryptone soya broth +2% 13 NaCl (TSB2) containing 200 micromolar ferric chloride 14 in a 2.5 l Erlenmeyer flask was inoculated with a 15 0.01% v/v inoculum of an 18h TSB2 culture of 16 Photobacterium damseale subsp. piscicida MT1415. The 17 culture was grown with shaking at 140 rpm until late 18 exponential growth phase (about 40 hours) at 24 °C. 19 The resulting culture was inactivated with formalin 20 21 (final concentration 0.2%), and protease was inactivated by adding Phenylmethylsulphonylfluoride 22 (PMSF) to a final concentration of 100 micromolar 23 from a 100 millimolar stock solution in isopropanol. 24 25 To prepare vaccines in which the expression of the 26 55KDa complex was completely inhibited. 27 Photobacterium damselae subsp. piscicida isolate 28 MT1415 was subcultureed twice for 18 hours in TSB2 29 30 containing 100 micromolar 2,2 dipyridyl, an iron chelator. This resulting completely iron-depleted 31 culture was used as the inoculum (0.5%v/v) for the 32

vaccine culture which was grown in 500 ml of TSB2

containing 100 micromolar 2,2 dipyridyl in a 2.5L

Erlenmeyer flask with shaking at 140 rpm until late

33

34

35

18

exponential growth phase (about 48 hours) at 24 C.

Absence of the 55Da complex from this preparation was

confirmed by western blot of outer membrane proteins

prepared from a duplicate culture (Refer to figure

7).

WO 01/10459

1 Claims

2

1. A composition comprising biological material derived from a culture of Photobacterium damselae, characterised in that the bacterial cells have been cultured in a culture medium containing excess iron over that which is required for' normal growth of the bacterium.

9

2. A composition as claimed in claim 1 wherein the biological material includes an outermembrane protein which is involved in entry into host cells, or invasin or adhesin and is expressed at levels higher than those in normal culture medium for use in the preparation of a vaccine.

16

3. A composition as claimed in claim 1 or claim 2
wherein an extracellular protein serologically
related to invasin is expressed at higher levels
than in normal culture.

21

4. A process for production of a vaccine comprising a step wherein bacterial cells of *Photobacterium damselae* are cultured in a culture medium containing sufficient iron such that the cells are loaded up with iron to the point where linear uptake of iron from the culture medium no longer occurs.

29

30 5. A process as claimed in claim 4 wherein the cells become saturated with iron.

32

33 6. A process as claimed in claim 4 or claim 5
34 wherein iron is supplied to the medium in the
35 form of a ferric salt.

WO 01/10459

1 2 7. A process as claimed in claim 4, 5 or 6 wherein 3 the bacteria are inactivated after culture. 4 8. A process as claimed in claim 7 wherein the 5 6 inactivation is carried out using a formaldehyde 7 composition. 8 9 9. A process as claimed in any of claims 4 to 8 10 wherein the products of the process are used for 11. the production of antibodies. 12 13 10. Use of cells, cell membranes or extracellular 14 15 products produced by the process as claimed in 16 any of claims 4 to 8 in the preparation of a vaccine for the prophylatic or therapeutic 17 treatment of fish from infection by bacteria. 18 19 20 11. Us of the composition as claimed in any of 21 claims 1 to 3 in the preparation of a vaccine for the prophylatic or therapeutic treatment of 22 fish from infection by bacteria. 23 24 25 12. A use as claimed in claim 10 or 11 wherein the vaccine includes a physiologically acceptable 26 27 carrier. 29 13. A use as claimed in claims 10 to 12 wherein the vaccine includes an adjuvant suitable for 30 enhancing immunological response. 31

28

32

33 14. A vaccine composition for the prophylactic 34 and/or therapeutic treatment of fish for infection by bacteria, particularly by the 35

1		organism Photobacterium damelae subsp. piscicida
2		wherein the composition comprises components
3		produced by a culture of the organism deposited
4		under Accession No 41062 at N.C.I.M.B. in
5		Aberdeen, United Kingdom on 4 August 2000.
6		
7	15.	Us of the bacterium deposited under Accession No
8		41062 at N.C.I.M.B. in Aberdeen, United Kingdom
9		on 4 August 2000 in the preparation of a
10		vaccine.
11		
12	16.	Antibodies produced by the process as claimed in
13	÷	claim 9.

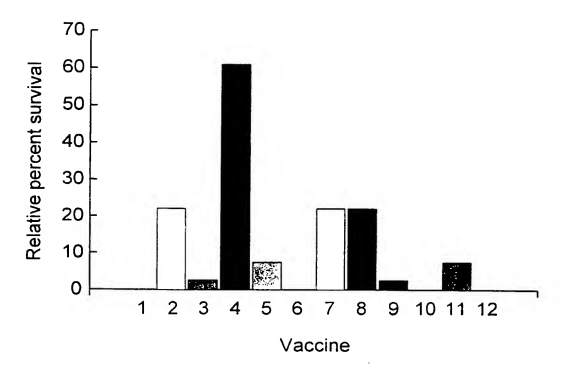
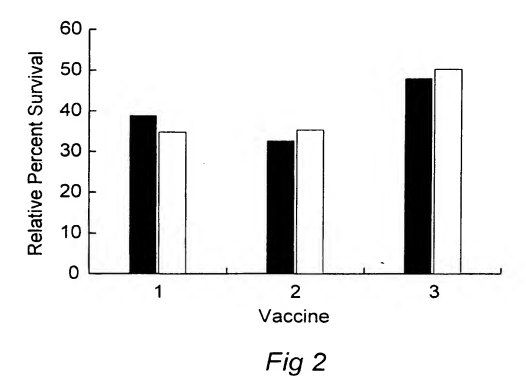
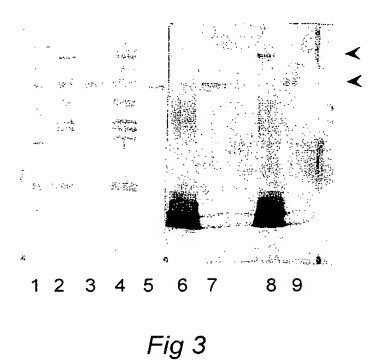
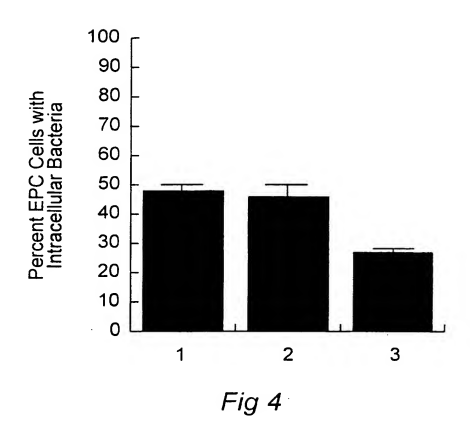


Fig 1



3/8





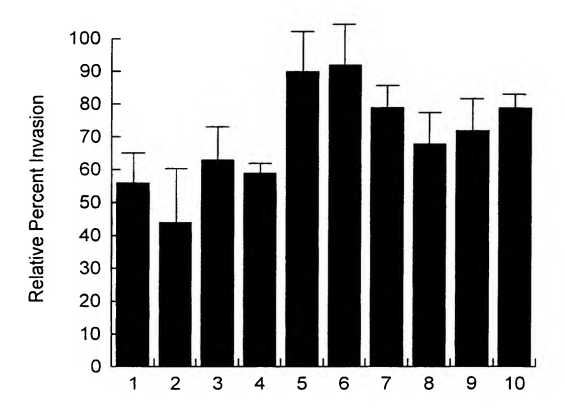


Fig 5

6/8

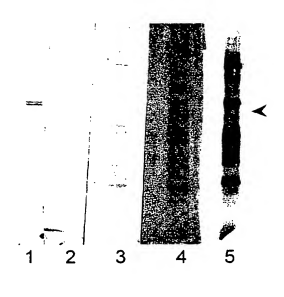


Fig 6

7/8



Fig 7

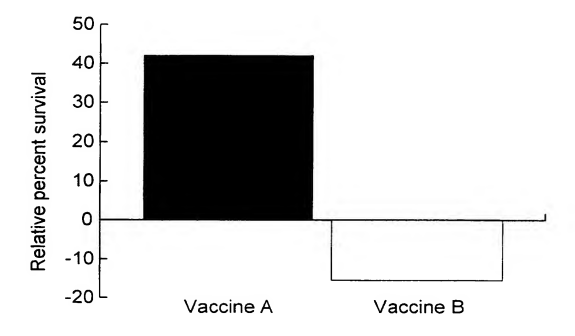


Fig 8

(19) World Intellectual Property Organization International Bureau



(43) International Publication Date 15 February 2001 (15.02.2001)

PCT

(10) International Publication Number WO 01/10459 A3

- (51) International Patent Classification7: A61K 39/106, 39/102, 39/02, C12N 1/20, C07K 16/12, A61P 31/04
- (21) International Application Number: PCT/GB00/02973
- (22) International Filing Date: 7 August 2000 (07.08.2000)
- (25) Filing Language:

English

(26) Publication Language:

English

(30) Priority Data:

9918591.0

7 August 1999 (07.08.1999) GB

9923030.2

30 September 1999 (30.09.1999) G

- (71) Applicant (for all designated States except US): AQUA HEALTH (EUROPE) LIMITED [GB/GB]; Enterprise House, Springkerse Business Park, Stirling FK7 7UF (GB).
- (72) Inventor; and
- (75) Inventor/Applicant (for US only): BARNES, Andrew, Cartner [GB/GB]; 54 Union Grove, Aberdeen AB10 6RX (GB).
- (74) Agent: MURGITROYD & COMPANY; 373 Scotland Street, Glasgow G5 8QA (GB).

- (81) Designated States (national): AE, AL, AM, AT, AU, AZ, BA, BB, BG, BR, BY, CA, CH, CN, CR, CU, CZ, DE, DK, DM, EE, ES, FI, GB, GD, GE, GH, GM, HR, HU, ID, IL, IN, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MA, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, TZ, UA, UG, US, UZ, VN, YU, ZA, ZW.
- (84) Designated States (regional): ARIPO patent (GH, GM, KE, LS, MW, MZ, SD, SL, SZ, TZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, CY, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, GW, ML, MR, NE, SN, TD, TG).

Published:

- With international search report.
- (88) Date of publication of the international search report: 10 May 2001

For two-letter codes and other abbreviations, refer to the "Guidance Notes on Codes and Abbreviations" appearing at the beginning of each regular issue of the PCT Gazette.

(54) Title: FISH VACCINE

(57) Abstract: A vaccine composition is provided for the prophylactic and/or therapeutic treatment of fish for infection by bacteria, particularly by the organism *Photobacterium damselae* subsp. *piscicida*. The composition comprises components produced by a culture of the organism. The culture is treated to kill the organism prior to use, preferably by treatment with formalin after components have been produced. Two of the components, involved in invasion of host cells and produced in high quantities when the culture is grown in medium containing excess iron, induce production of antibodies on injection into fish which prevent the entry of the organism into fish cells. These antibodies protect the fish from infection by the organism.

INTERNATIONAL SEARCH REPORT

Interr. Ial Application No

PCT/GB 00/02973

A. CLASS	IFICATION OF SUBJECT MATTER	4						
IPC 7	A61K39/106 A61K39/102 A61K39	7/02 C12N1/20 C	07K16/12					
	A61P31/04		-					
	o International Patent Classification (IPC) or to both national class	arication and IPC	•					
	SEARCHED	nation of the state of the stat						
IPC 7	ocumentation searched (classification system followed by classific A61K C12N C07K	cation symbols)	÷					
	• • • • • • • • • • • • • • • • • • • •							
		 						
Documenta	tion searched other than minimum documentation to the extent the	at such documents are included in the fie	lds searched					
Electronic d	ata base consulted during the international search (name of data	base and where practical search terms	used)					
		•						
WFI Da	ta, EPO-Internal, PAJ, BIOSIS, MED	LINE, CHEM ABS Data						
C DOCUM	ENTS CONSIDERED TO BE RELEVANT	· · · · · · · · · · · · · · · · · · ·						
Category *	Citation of document, with indication, where appropriate, of the	misuset soosses	Bolovenia del N					
Calegory	Chanon of document, with indication, where appropriate, of the	relevani passages	Relevant to claim No.					
X	WO 96 12734 A (EWOS AKTIEBOLAG)		1,2,4-16					
	2 May 1996 (1996-05-02)	•						
	page 2, line 20 -page 3, line 5	; claims						
	1,2,4,5,9,17,18,20-27							
			1					
,								
			.v.					
1								
l								
			l l					
Furth	er documents are listed in the continuation of box C.	Patent family members are list	sted in annex.					
° Special cat	egories of cited documents:	"T" later document published after the	international filing date					
A docume	nt defining the general state of the art which is not	or priority date and not in conflict	with the application but					
	ered to be of particular relevance	cited to understand the principle of invention						
filing da	ocument but published on or after the international ste	"X" document of particular refevance; t cannot be considered novel or can						
"L" documer which is	document is taken alone							
citation	he claimed invention n inventive step when the							
O docume other m	r more other such docu- ovious to a person skilled							
P documer								
	an the priority date claimed	*&* document member of the same pat						
Date of the a	ctual completion of the international search	Date of mailing of the international	search report					
	- Tanuary 2001	05 (00 (000)						
30	January 2001	05/02/2001						
Name and m	ailing address of the ISA	Authorized officer						
European Patent Office, P.B. 5818 Patentlaan 2 NL - 2280 HV Rijswijk								
	I							

INTERNATIONAL SEARCH REPORT

information on patent family members

Intern nal Application No

						FCI/GB	00/029/3	
cite	atent document d in search repor	t	Publication date	þ	atent family nember(s)	,	Publication date	
WO	9612734	Α	02-05-1996	AU	38210	95 A	15-05-1996	
							•	

Form PCT/ISA/210 (patent family annex) (July 1992)